Université de BORDEAUX

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Biopython: Multiple alignments





Introduction

- → A collection of multiple sequences which have been aligned together
- → Insertion of gap characters, and addition of leading or trailing gaps
 - > such that all the sequence strings are the same length.
- → Can be regarded as a matrix of letters, where each row is held as a SeqRecord object internally
- → Introduce :
 - MultipleSeqAlignment object which holds this kind of data
 - Bio.AlignIO module for reading and writing them as various file formats (following the design of the Bio.SeqIO module)



Parsing or Reading Sequence Alignments

- → Bio.AlignIO.read() and Bio.AlignIO.parse()
- Using Bio.AlignIO.parse() will return an iterator which gives MultipleSeqAlignment objects
- Bio.AlignIO.read() function which returns a single MultipleSeqAlignment object
- → First argument is a handle to read the data from, typically an open file, or a filename
- → Second argument is a lower case string specifying the alignment format. (see http://biopython.org/wiki/AlignIO)
- → Optional seq_count argument for dealing with ambiguous file formats
- Optional Alphabet argument allowing you to specify the expected alphabet.



Single Alignments

consider the following annotation rich protein alignment in the PFAM or Stockholm file

```
# STOCKHOLM 1.0
#=GS COATB BPIKE/30-81 AC P03620.1
#=GS COATB BPIKE/30-81 DR PDB; 1ifl : 1-52;
#=GS Q9T0Q8 BPIKE/1-52 AC Q9T0Q8.1
#=GS COATB BPI22/32-83 AC P15416.1
#=GS COATB BPM13/24-72 AC P69541.1
#=GS COATB BPM13/24-72 DR PDB; 2cpb; 1-49;
#=GS COATB BPM13/24-72 DR PDB; 2cps : 1-49;
#=GS COATB BPZJ2/1-49 AC P03618.1
#=GS Q9T0Q9 BPFD/1-49 AC Q9T0Q9.1
#=GS Q9T0Q9 BPFD/1-49 DR PDB; 1nh4 A; 1-49;
#=GS COATB BPIF1/22-73 AC P03619.2
#=GS COATB BPIF1/22-73 DR PDB; 1ifk; 1-50;
COATB BPIKE/30-81 AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA
Q9T0Q8 BPIKE/1-52 AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA
COATB BPI22/32-83 DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA
COATB BPZJ2/1-49 AEGDDP...AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFASKA
Q9T0Q9 BPFD/1-49 AEGDDP...AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA
COATB BPIF1/22-73 FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA
#=GC seg cons AEssss...AptAhDSLpspAT-hlu.sWshVsslVsAslulKLFKKFsSK
```

Single Alignments (2)

>>> from Bio import AlignIO

```
>>> from Bio import AlignIO
>>> alignment = AlignIO.read("PF05371_seed.sth", "stockholm")

>>> print(alignment)
SingleLetterAlphabet() alignment with 7 rows and 52 columns
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRL...SKA COATB_BPIKE/30-81
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKL...SRA Q9T0Q8_BPIKE/1-52
DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRL...SKA COATB_BPI22/32-83
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPM13/24-72
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPZJ2/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA Q9T0Q9_BPFD/1-49
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKL...SRA COATB_BPIF1/22-73
```

```
>>> alignment = AlignIO.read("PF05371_seed.sth", "stockholm")
>>> print("Alignment length %i" % alignment.get_alignment_length())
Alignment length 52
>>> for record in alignment:
... print("%s - %s" % (record.seq, record.id))
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA - COATB_BPIKE/30-81
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA - Q9T0Q8_BPIKE/1-52
DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA - COATB_BPI22/32-83
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA - COATB_BPM13/24-72
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA - Q9T0Q9_BPFD/1-49
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA - COATB_BPIF1/22-73
```



Single Alignments (3)

→ Include database cross-references to the PDB and the associate known secondary structure

```
>>> for record in alignment:
... if record.dbxrefs:
... print("%s %s" % (record.id, record.dbxrefs))
COATB BPIKE/30-81 ['PDB; 1ifl; 1-52;']
COATB BPM13/24-72 ['PDB; 2cpb; 1-49;', 'PDB; 2cps; 1-49;']
Q9T0Q9 BPFD/1-49 ['PDB; 1nh4 A; 1-49;']
COATB BPIF1/22-73 ['PDB; 1ifk; 1-50;']
>>> for record in alignment:
... print(record)
>COATB BPIKE/30-81
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA
>Q9T0Q8 BPIKE/1-52
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA
>COATB_BPI22/32-83
DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA
>COATB BPM13/24-72
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA
>COATB BPZJ2/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFASKA
>Q9T0Q9 BPFD/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA
>COATB BPIF1/22-73
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA
```



Single Alignments (4)

→ download and save this as file "PF05371 seed.fa"

```
from Bio import AlignIO
alignment = AlignIO.read("PF05371_seed.faa", "fasta")
print(alignment)
```

→ no annotation nor database cross-references because these are not included in the FASTA file format



Multiple alignments (1)

→ Suppose you have a small alignment in PHYLIP format

5 6 Alpha AACAAC Beta AACCCC Gamma ACCAAC Delta CCACCA Epsilon CCAAAC

- → To bootstrap a phylogenetic tree using the PHYLIP tools:
 - create a set of many resampled alignments using the tool bootseq
- → This would give output (abbreviated for conciseness):

56 Alpha AAACCA Beta AAACCC Gamma ACCCCA Delta CCCAAC **Epsilon CCCAAA** 56 Alpha AAACAA Beta AAACCC Gamma ACCCAA Delta CCCACC **Epsilon CCCAAA** 56 Alpha AAAAAC Beta AAACCC Gamma AACAAC Delta CCCCCA

Multiple alignments (2)

→ If you wanted to read this in using Bio.AlignIO you could use

```
from Bio import AlignIO
alignments = AlignIO.parse("resampled.phy", "phylip")
for alignment in alignments:
print(alignment)
print("")
```

→ abbreviated for display:

```
SingleLetterAlphabet() alignment with 5 rows and 6 columns
AAACCA Alpha
AAACCC Beta
ACCCCA Gamma
CCCAAC Delta
CCCAAA Epsilon
SingleLetterAlphabet() alignment with 5 rows and 6 columns
AAACAA Alpha
AAACCC Beta
ACCCAA Gamma
CCCACC Delta
CCCAAA Epsilon
SingleLetterAlphabet() alignment with 5 rows and 6 columns
AAAACC Alpha
ACCCCC Beta
AAAACC Gamma
CCCCAA Delta
```



CAAACC Epsilon

Multiple alignments (3)

- → As with the function Bio.SeqIO.parse(), using Bio.AlignIO.parse() returns an iterator.
- → to keep all the alignments in memory at once
- → Turn the iterator into a list

```
from Bio import AlignIO
alignments = list(AlignIO.parse("resampled.phy", "phylip"))
last_align = alignments[-1]
first_align = alignments[0]
```



Ambiguous alignments (&)

→ Many alignment file formats can explicitly store more than one alignment,

>Alpha
ACTACGACTAGCTCAG--G
>Beta
ACTACCGCTAGCTCAGAAG
>Gamma
ACTACGGCTAGCACAGAAG
>Alpha
ACTACGACTAGCTCAGG->Beta
ACTACCGCTAGCTCAGAAG
>Gamma
ACTACCGCTAGCTCAGAAG

→ What about this next example?

>Alpha
ACTACGACTAGCTCAG--G
>Beta
ACTACCGCTAGCTCAGAAG
>Alpha
ACTACGACTAGCTCAGG->Gamma
ACTACGGCTAGCACAGAAG
>Alpha
ACTACGACTAGCTCAGG->Delta
ACTACGGCTAGCACAGAAG

→ To interpret these FASTA examples as several separate alignments, we can use Bio.AlignIO.parse() with the optional seq_count argument which specifies how many sequences are expected in each alignment (3,2,2 in the previous example

```
for alignment in AlignIO.parse(handle, "fasta", seq_count=2):
print("Alignment length %i" % alignment.get_alignment_length())
for record in alignment:
print("%s - %s" % (record.seq, record.id))
print("")
```

```
Alignment length 19
ACTACGACTAGCTCAG--G - Alpha
ACTACCGCTAGCTCAGAAG - XXX
Alignment length 17
ACTACGACTAGCTCAGG - Alpha
ACTACGGCAAGCACAGG - YYY
Alignment length 21
--ACTACGAC--TAGCTCAGG - Alpha
GGACTACGACAATAGCTCAGG - ZZZ
```



Writing alignments (1)

→ Bio.AlignIO.write() taking three arguments:

- Some MultipleSeqAlignment objects (or for backwards compatibility the Obsolete Alignment objects),
- > a handle or filename to write to
- a sequence format

```
from Bio.Alphabet import generic dna
from Bio.Seq import Seq
from Bio.SegRecord import SegRecord
from Bio.Align import MultipleSeqAlignment
align1 = MultipleSegAlignment([
SeqRecord(Seq("ACTGCTAGCTAG", generic dna), id="Alpha"),
SegRecord(Seg("ACT-CTAGCTAG", generic dna), id="Beta"),
SegRecord(Seg("ACTGCTAGDTAG", generic dna), id="Gamma"),
align2 = MultipleSegAlignment([
SeqRecord(Seq("GTCAGC-AG", generic dna), id="Delta"),
SeqRecord(Seq("GACAGCTAG", generic dna), id="Epsilon"),
SegRecord(Seg("GTCAGCTAG", generic dna), id="Zeta"),
align3 = MultipleSeqAlignment([
SeqRecord(Seq("ACTAGTACAGCTG", generic dna), id="Eta"),
SegRecord(Seg("ACTAGTACAGCT-", generic dna), id="Theta"),
SegRecord(Seg("-CTACTACAGGTG", generic dna), id="lota"),
])
my alignments = [align1, align2, align3]
```



→ Now we have a list of Alignment objects, we'll write them to a PHYLIP format file:

from Bio import AlignIO AlignIO.write(my_alignments, "my_example.phy", "phylip")

→ open this file in your favourite text editor it should look like this:

3 12
Alpha ACTGCTAGCT AG
Beta ACT-CTAGCT AG
Gamma ACTGCTAGDT AG
3 9
Delta GTCAGC-AG
Epislon GACAGCTAG
Zeta GTCAGCTAG
3 13
Eta ACTAGTACAG CTG
Theta ACTAGTACAG GTG



Converting between sequence alignment file formats (1)

→ For this example, we'll load the PFAM/Stockholm format file used earlier and save it as a Clustal W format file:

```
from Bio import AlignIO count = AlignIO.convert("PF05371_seed.sth", "stockholm", "PF05371_seed.aln", "clustal") print("Converted %i alignments" % count)
```

→ we gave it the alignment iterator returned by Bio.AlignIO.parse()

```
from Bio import AlignIO alignments = AlignIO.parse("PF05371_seed.sth", "stockholm") count = AlignIO.write(alignments, "PF05371_seed.aln", "clustal") print("Converted %i alignments" % count)
```

→ Or, using Bio.AlignIO.parse() and Bio.AlignIO.write()

```
from Bio import AlignIO alignment = AlignIO.read("PF05371_seed.sth", "stockholm") AlignIO.write([alignment], "PF05371_seed.aln", "clustal")
```



Converting between sequence alignment file formats (2)

→ you should end up with the same new Clustal W format file "PF05371 seed.aln" with the following content

```
CLUSTAL X (1.81) multiple sequence alignment
COATB_BPIKE/30-81 AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSS
Q9T0Q8_BPIKE/1-52 AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVS
COATB_BPI22/32-83 DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSS
COATB_BPM13/24-72 AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTS
COATB_BPZJ2/1-49 AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFAS
Q9T0Q9_BPFD/1-49 AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTS
COATB_BPIF1/22-73 FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVS
COATB_BPIKE/30-81 KA
Q9T0Q8_BPIKE/1-52 RA
COATB_BPI22/32-83 KA
COATB_BPM13/24-72 KA
COATB_BPZJ2/1-49 KA
Q9T0Q9_BPFD/1-49 KA
COATB_BPI1/22-73 RA
```

→ you could make a PHYLIP format le which we'll name "PF05371 seed.phy"

```
from Bio import AlignIO
AlignIO.convert("PF05371_seed.sth", "stockholm", "PF05371_seed.phy", "phylip")
```



Converting between sequence alignment file formats (3)

COATB_BPIK AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIRLFKKFSS Q9T0Q8_BPI AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIKLFKKFVS COATB_BPI2 DGTSTATSYA TEAMNSLKTQ ATDLIDQTWP VVTSVAVAGL AIRLFKKFSS COATB_BPM1 AEGDDP---A KAAFNSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS COATB_BPZJ AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFAS Q9T0Q9_BPF AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS COATB_BPIF FAADDATSQA KAAFDSLTAQ ATEMSGYAWA LVVLVVGATV GIKLFKKFVS KA RA

→ Big handicaps of the original PHYLIP alignment file format is that the sequence identifiers are strictly truncated at ten characters.

```
from Bio import AlignIO
AlignIO.convert("PF05371_seed.sth", "stockholm", "PF05371_seed.phy", "phylip-relaxed")
7 52
COATB_BPIKE/30-81 AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIRLFKKFSS
Q9T0Q8_BPIKE/1-52 AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIKLFKKFVS
COATB_BPI22/32-83 DGTSTATSYA TEAMNSLKTQ ATDLIDQTWP VVTSVAVAGL AIRLFKKFSS
COATB_BPM13/24-72 AEGDDP---A KAAFNSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS
COATB_BPZJ2/1-49 AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFAS
Q9T0Q9_BPFD/1-49 AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS
COATB_BPIF1/22-73 FAADDATSQA KAAFDSLTAQ ATEMSGYAWA LVVLVVGATV GIKLFKKFVS
KA
RA
....
```



Converting between sequence alignment file formats (4)

→ have to work with the original strict PHYLIP format, then you may need to compress the identifiers somehow or assign your own names or numbering system.

```
from Bio import AlignIO
alignment = AlignIO.read("PF05371_seed.sth", "stockholm")
name_mapping = {}
for i, record in enumerate(alignment):
name_mapping[i] = record.id
record.id = "seq%i" % i
print(name_mapping)
AlignIO.write([alignment], "PF05371_seed.phy", "phylip")
```

use a Python dictionary to record a simple mapping from the new sequence system to the original identifier:

```
{0: 'COATB BPIKE/30-81', 1: 'Q9T0Q8 BPIKE/1-52', 2: 'COATB BPI22/32-83', ...}
```

```
7 52
seq0 AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIRLFKKFSS
seq1 AEPNAATNYA TEAMDSLKTQ AIDLISQTWP VVTTVVVAGL VIKLFKKFVS
seq2 DGTSTATSYA TEAMNSLKTQ ATDLIDQTWP VVTSVAVAGL AIRLFKKFSS
seq3 AEGDDP---A KAAFNSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS
seq4 AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFAS
seq5 AEGDDP---A KAAFDSLQAS ATEYIGYAWA MVVVIVGATI GIKLFKKFTS
seq6 FAADDATSQA KAAFDSLTAQ ATEMSGYAWA LVVLVVGATV GIKLFKKFVS
KA
```



Getting your alignment in formatted string (1)

→ Take advantage of the alignment object's format() method

```
from Bio import AlignIO
alignment = AlignIO.read("PF05371_seed.sth", "stockholm")
print(alignment.format("clustal"))
```

- → The SeqRecord object has a similar method using output formats supported by Bio.SeqIO
- → The format() method is using the StringIO string based handle and calling Bio.AlignIO.write()

```
from Bio import AlignIO
from StringIO import StringIO
alignments = AlignIO.parse("PF05371_seed.sth", "stockholm")
out_handle = StringIO()
AlignIO.write(alignments, out_handle, "clustal")
clustal_data = out_handle.getvalue()
print(clustal_data)
```



Slicing alignments (1)

→ Alignment objects act like a Python list of SeqRecord objects (the rows)

```
>>> from Bio import AlignIO
>>> alignment = AlignIO.read("PF05371_seed.sth", "stockholm")
>>> print("Number of rows: %i" % len(alignment))
Number of rows: 7
>>> for record in alignment:
... print("%s - %s" % (record.seq, record.id))
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA - Q9T0Q8_BPIKE/1-52
DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA - COATB_BPI22/32-83
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA - COATB_BPM13/24-72
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA - COATB_BPZJ2/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA - Q9T0Q9_BPFD/1-49
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA - COATB_BPIF1/22-73
```

→ Suppose you have a small alignment in PHYLIP format

```
>>> print(alignment)
SingleLetterAlphabet() alignment with 7 rows and 52 columns
AEPNAATNYATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKL...SRA Q9T0Q8_BPIKE/1-52
DGTSTATSYATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRL...SKA COATB_BPI22/32-83
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPM13/24-72
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPZJ2/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA Q9T0Q9_BPFD/1-49
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKL...SRA COATB_BPIF1/22-73
>>> print(alignment[3:7])
SingleLetterAlphabet() alignment with 4 rows and 52 columns
AEGDDP---AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPM13/24-72
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA COATB_BPZJ2/1-49
AEGDDP---AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKL...SKA Q9T0Q9_BPFD/1-49
FAADDATSQAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKL...SRA COATB_BPIF1/22-73
```

Alignments as array (1)

→ If you wanted to select by column?

```
>>> print(alignment[2, 6])
T
```

→ Using two integer indices pulls out a single letter

```
>>> print(alignment[2].seq[6])
```

→ pull out a single column as a string like this

```
>>> print(alignment[:, 6])
TTT---T
```

→ Select a range of columns

```
>>> print(alignment[3:6, :6])
SingleLetterAlphabet() alignment with 3 rows and 6 columns
AEGDDP COATB_BPM13/24-72
AEGDDP COATB_BPZJ2/1-49
AEGDDP Q9T0Q9_BPFD/1-49
```

→ Leaving the first index as means take all the rows:

```
>>> print(alignment[:, :6])
SingleLetterAlphabet() alignment with 7 rows and 6 columns
AEPNAA COATB_BPIKE/30-81
AEPNAA Q9T0Q8_BPIKE/1-52
.....
```

Slicing alignments (1)

→ columns 7, 8 and 9 which are gaps in three of the seven sequences

```
>>> print(alignment[:, 6:9])
SingleLetterAlphabet() alignment with 7 rows and 3 columns
TNY COATB_BPIKE/30-81
TNY Q9T0Q8_BPIKE/1-52
TSY COATB_BPI22/32-83
--- COATB_BPM13/24-72
--- COATB_BPZJ2/1-49
--- Q9T0Q9_BPFD/1-49
TSQ COATB_BPIF1/22-73
```

→ you can slice to get everything after the ninth column

```
>>> print(alignment[:, 9:])
SingleLetterAlphabet() alignment with 7 rows and 43 columns
ATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA COATB_BPIKE/30-81
ATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA Q9T0Q8_BPIKE/1-52
ATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA COATB_BPI22/32-83
AKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA COATB_BPM13/24-72
AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFASKA COATB_BPZJ2/1-49
AKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA Q9T0Q9_BPFD/1-49
AKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA COATB_BPIF1/22-73
```



Slicing alignments (1)

→ Addition of alignment objects works by column

```
>>> edited = alignment[:, :6] + alignment[:, 9:]
>>> print(edited)
SingleLetterAlphabet() alignment with 7 rows and 49 columns
AEPNAAATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA COATB_BPIKE/30-81
AEPNAAATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA Q9T0Q8_BPIKE/1-52
DGTSTAATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA COATB_BPI22/32-83
AEGDDPAKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA COATB_BPM13/24-72
AEGDDPAKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFASKA COATB_BPZJ2/1-49
AEGDDPAKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA Q9T0Q9_BPFD/1-49
FAADDAAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA COATB_BPIF1/22-73
```

→ Combine alignments for several different genes into a meta-alignment

```
>>> edited.sort()
>>> print(edited)
SingleLetterAlphabet() alignment with 7 rows and 49 columns
DGTSTAATEAMNSLKTQATDLIDQTWPVVTSVAVAGLAIRLFKKFSSKA COATB_BPI22/32-83
FAADDAAKAAFDSLTAQATEMSGYAWALVVLVVGATVGIKLFKKFVSRA COATB_BPIF1/22-73
AEPNAAATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIRLFKKFSSKA COATB_BPIKE/30-81
AEGDDPAKAAFNSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA COATB_BPM13/24-72
AEGDDPAKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFASKA COATB_BPZJ2/1-49
AEPNAAATEAMDSLKTQAIDLISQTWPVVTTVVVAGLVIKLFKKFVSRA Q9T0Q8_BPIKE/1-52
AEGDDPAKAAFDSLQASATEYIGYAWAMVVVIVGATIGIKLFKKFTSKA Q9T0Q9_BPFD/1-49
```

Only add two alignments together if they have the same number of rows

Alignments as array (1)

→ Suppose you have a small alignment in PHYLIP format

```
>>> import numpy as np
>>> from Bio import AlignIO
>>> alignment = AlignIO.read("PF05371_seed.sth", "stockholm")
>>> align_array = np.array([list(rec) for rec in alignment], np.character)
>>> print("Array shape %i by %i" % align_array.shape)
Array shape 7 by 52
```

→ Tell NumPy to store the array by column (as in Fortran) rather then its default of by row (as in C)

```
>>> align_array = np.array([list(rec) for rec in alignment], np.character, order="F")
```

→ Leave the original Biopython alignment object and the NumPy array in memory as separate objects



Alignment tools (1)

- Prepare an input file of your unaligned sequences, typically this will be a FASTA le which you might create using Bio.SeqIO
- Call the command line tool to process this input file, typically via one of Biopython's command line wrappers (which we'll discuss here).
- Read the output from the tool, i.e. your aligned sequences, typically using Bio.AlignIO

```
>>> import Bio.Align.Applications
>>> dir(Bio.Align.Applications)
['ClustalwCommandline', 'DialignCommandline', 'MafftCommandline', 'MuscleCommandline',
'PrankCommandline', 'ProbconsCommandline', 'TCoffeeCommandline' ...]
```



ClustalW (1)

 ClustalW is a popular command line tool for multiple sequence alignment

```
>>> from Bio.Align.Applications import ClustalwCommandline
>>> help(ClustalwCommandline)
....
```

→ By default, ClustalW will generate an alignment and guide tree file with names based on the input FASTA file.

```
>>> from Bio.Align.Applications import ClustalwCommandline
>>> cline = ClustalwCommandline("clustalw2", infile="opuntia.fasta")
>>> print(cline)
clustalw2 -infile=opuntia.fasta
```

```
>>> import os
>>> from Bio.Align.Applications import ClustalwCommandline
>>> clustalw_exe = r"C:\Program Files\new clustal\clustalw2.exe"
>>> clustalw_cline = ClustalwCommandline(clustalw_exe, infile="opuntia.fasta")
>>> assert os.path.isfile(clustalw_exe), "Clustal W executable missing"
>>> stdout, stderr = clustalw_cline()
```



ClustalW (2)

ClustalW is a popular command line tool for multiple sequence alignment

```
>>> from Bio import AlignIO
>>> align = AlignIO.read("opuntia.aln", "clustal")
>>> print(align)
SingleLetterAlphabet() alignment with 7 rows and 906 columns
TATACATTAAAGAAGGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA gi|6273285|gb|AF191659.1|AF191
TATACATTAAAGAAGGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA qi|6273284|qb|AF191658.1|AF191
TATACATTAAAGAAGGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA gi|6273287|gb|AF191661.1|AF191
TATACATAAAAGAAGGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA gil6273286|gb|AF191660.1|AF191
TATACATTAAAGGAGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA qil6273290|qb|AF191664.1|AF191
TATACATTAAAGGAGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA gi|6273289|gb|AF191663.1|AF191
TATACATTAAAGGAGGGGATGCGGATAAATGGAAAGGCGAAAG...AGA gil6273291|gb|AF191665.1|AF191
>>> from Bio import Phylo
>>> tree = Phylo.read("opuntia.dnd", "newick")
>>> Phylo.draw ascii(tree)
               gi|6273291|gb|AF191665.1|AF191665
        gi|6273290|gb|AF191664.1|AF191664
       gi|6273289|gb|AF191663.1|AF191663
                   gi|6273287|gb|AF191661.1|AF191661
           gi|6273286|gb|AF191660.1|AF191660
   gi|6273285|gb|AF191659.1|AF191659
| gi|6273284|gb|AF191658.1|AF191658
<BLANKLINE>
```

EMBOSS needle and water (1)

- → The EMBOSS suite includes the water and needle tools for Smith-Waterman algorithm local alignment, and Needleman-Wunsch global alignment
- → Suppose you want to do a global pairwise alignment between two sequences, prepared in FASTA format (alpha.faa and beta.faa)

>HBA HUMAN

MVLSPADKTNVKAAWGKVGAHAGEYGAEALERMFLSFPTTKTYFPHFDLSHGSAQVKGHG KKVADALTNAVAHVDDMPNALSALSDLHAHKLRVDPVNFKLLSHCLLVTLAAHLPAEFTP AVHASLDKFLASVSTVLTSKYR

>HBB HUMAN

MVHLTPEEKSAVTALWGKVNVDEVGGEALGRLLVVYPWTQRFFESFGDLSTPDAVMGNPK VKAHGKKVLGAFSDGLAHLDNLKGTFATLSELHCDKLHVDPENFRLLGNVLVCVLAHHFG KEFTPPVQAAYQKVVAGVANALAHKYH

→ Let's start by creating a complete needle command line object in one go:

```
>>> from Bio.Emboss.Applications import NeedleCommandline
>>> needle_cline = NeedleCommandline(asequence="alpha.faa", bsequence="beta.faa", gapopen=10, gapextend=0.5, outfile="needle.txt")
>>> print(needle_cline)
needle -outfile=needle.txt -asequence=alpha.faa -bsequence=beta.faa -gapopen=10 -gapextend=0.5
```



EMBOSS needle and water (2)

→ To avoid "command not found" exception, you can either update your PATH setting, or simply tell Biopython the full path to the tool, for example:

```
>>> from Bio.Emboss.Applications import NeedleCommandline
>>> needle_cline = NeedleCommandline(r"C:\EMBOSS\needle.exe",
... asequence="alpha.faa", bsequence="beta.faa",
... gapopen=10, gapextend=0.5, outfile="needle.txt")
```

→ Note that you can also specify (or change or look at) the settings like this:

```
>>> from Bio.Emboss.Applications import NeedleCommandline
>>> needle_cline = NeedleCommandline()
>>> needle_cline.asequence="alpha.faa"
>>> needle_cline.bsequence="beta.faa"
>>> needle_cline.gapopen=10
>>> needle_cline.gapextend=0.5
>>> needle_cline.outfile="needle.txt"
>>> print(needle_cline)
needle - outfile=needle.txt - asequence=alpha.faa - bsequence=beta.faa - gapopen=10 - gapextend=0.5
>>> print(needle_cline.outfile)
needle.txt
```



EMBOSS needle and water (3)

→ To avoid "command not found" exception, you can either update your PATH setting, or simply tell Biopython the full path to the tool, for example:

```
>>> from Bio.Emboss.Applications import NeedleCommandline
>>> needle_cline = NeedleCommandline(r"C:\EMBOSS\needle.exe",
... asequence="alpha.faa", bsequence="beta.faa",
... gapopen=10, gapextend=0.5, outfile="needle.txt")
```

→ Note that you can also specify (or change or look at) the settings like this:

```
>>> from Bio.Emboss.Applications import NeedleCommandline
>>> needle_cline = NeedleCommandline()
>>> needle_cline.asequence="alpha.faa"
>>> needle_cline.bsequence="beta.faa"
>>> needle_cline.gapopen=10
>>> needle_cline.gapextend=0.5
>>> needle_cline.outfile="needle.txt"
>>> print(needle_cline)
needle -outfile=needle.txt -asequence=alpha.faa -bsequence=beta.faa -gapopen=10 -gapextend=0.5
>>> print(needle_cline.outfile)
needle.txt
```



Multiple alignments (3)

→ Suppose you have a small alignment in PHYLIP format

5 6 Alpha AACAAC Beta AACCCC Gamma ACCAAC Delta CCACCA Epsilon CCAAAC

→ Suppose you have a small alignment in PHYLIP format

5 6 Alpha AACAAC Beta AACCCC Gamma ACCAAC Delta CCACCA Epsilon CCAAAC

